REMARKS

As of the Preliminary Amendment filed on February 28, 2003, claims 1-14 were pending in the present application. Claims 2, 4, and 11 are canceled herein. Claims 3, 5, 6, and 13 are amended. Claims 15-27 are added. No new matter is inserted into the application.

Status of the Application

The Office Action dated February 25, 2003 presents the examination of claims 1-6, 10, and 11, which remain rejected. The mailing date of the Office Action precedes the date of the Preliminary Amendment filed on February 28, 2003. Thus, the Examiner did not consider the Preliminary Amendment in connection with the outstanding Office Action, as noted in the Communication from the Examiner dated March 14, 2003. In the Communication, the Examiner states that the Preliminary Amendment does not address all issues of record in the outstanding Office Applicants respectfully submit that the instant Reply addresses all remaining issues of the Office Action and is therefore a complete response. Upon entry of this Reply, claims 1, 3, 5-10, and 12-27 should be pending.

Rejection under 35 U.S.C. § 112, first paragraph, new matter

The Examiner maintains the rejection of claim 1 under 35 U.S.C. § 112, first paragraph for allegedly containing new matter. Applicants respectfully traverse. Reconsideration of the claim and withdrawal of the instant rejection are respectfully requested.

The Examiner asserts that there is no support in the specification for a mutant α -amylase comprising an amino acid sequence which is at least 70% homologous to SEQ ID NO:1, as recited in claim 1. Applicants respectfully disagree. Support for a mutant α -amylase comprising an amino acid sequence which is at least 70% homologous to SEQ ID NO:1 is found throughout the specification, such as on page 3, lines 9-10 and 16-17, page 4, lines 23-25, and in the abstract. Furthermore, SEQ ID NO:2 is a liquefying alkaline α -amylase having 66.9% identity to SEQ ID NO:1. In this regard, the Examiner's attention is drawn to Exhibit 1 (attached hereto) which provides a sequence alignment between SEQ ID NO:1 and SEQ ID NO:2, using the matrix file BLOSUM50. SEQ ID NO:2 is referred to in the specification at least on page 7, line 21 to page 8, line 2.

For this reason, the rejection is improper. Withdrawal thereof is respectfully requested.

Rejection under 35 U.S.C. § 112, first paragraph, written description

The Examiner maintains the rejection of claims 1, 5 and 6 under 35 U.S.C. § 112, first paragraph for allegedly not being described in the specification. Applicants respectfully traverse. Reconsideration of the claims and withdrawal of the instant rejection are respectfully requested.

Claim 1

The Examiner maintains that the claim 1 encompasses a protein variant having an untold number of mutations. Applicants respectfully disagree.

Claim 1 is directed to a mutant α -amylase obtained by making a substitution or deletion of at least one amino acid residue of specific positions in SEQ ID NO:1, or by making a substitution or deletion of at least one amino acid residue corresponding to the above-mentioned amino acid residue in a sequence having at least 70% homology to SEQ ID NO:1, wherein said at least one amino acid residue is selected from the group consisting of: the 11th Tyr, 16th Glu, 49th Asn, 84th Glu, 144th Ser, 167th Gln, 169th Tyr, 178th Ala, 188th Glu, 190th Asn, 205th His and 209th Gln, and said mutant α -amylase possesses increased heat resistance and maintains resistance to chelating agents when compared to SEQ ID NO:1, and said mutant α -amylase comprises an

amino acid sequence which is at least 70% homologous to SEQ ID NO:1. In other words, the mutant α -amylase must (1) be derived from an amino acid sequence corresponding to SEQ ID NO:1 or an amino acid sequence 70% homologous to SEQ ID NO:1, (2) have a substitution or deletion of the 11th Tyr, 16th Glu, 49th Asn, 84th Glu, 144th Ser, 167th Gln, 169th Tyr, 178th Ala, 188th Glu, 190th Asn, 205th His, or the 209th Gln, (3) possess the specific functions of increased heat resistance and resistance to chelating agents, and (4) have a resulting amino acid sequence which is at least 70% homologous to SEQ ID NO:1. Thus, the mutant α -amylase mutant is defined by several limitations in the claim.

The specification provides several examples of mutant α -amylases according to the present invention. As noted above, SEQ ID NO:2 is a liquefying alkaline α -amylase having 66.9% identity to SEQ ID NO:1 (see Exhibit 1). SEQ ID NO:4 is a liquefying alkaline α -amylase having 96.5% identity to SEQ ID NO:1. In this regard, the Examiner's attention is drawn to Exhibit 2 (attached hereto) which provides a sequence alignment between SEQ ID NO:1 and SEQ ID NO:4, using the matrix file BLOSUM50. SEQ ID NO:4 is referred to in the specification at least on page 5, lines 19-27. Other examples, although not listed by a particular SEQ ID, are described in Examples 5-10, pages 22-27 of the specification.

For the Examiner's convenience, the particular examples disclosed in the specification are listed in the following table:

MUTANT α-AMYLASE	SUPPORT IN
	SPECIFICATION
Y11F	Page 23
N49S	Page 23
E84Q	Page 23
S144P	Page 23
Q167E	Page 23
Y169K	Page 23
A178Q	Page 23
E188D	Page 23
N190F	Page 23
Q209V	Page 23
Q167E/Y169K	Page 24
N190F/Q209V	Page 24
Q167E/Y169K/N190F/Q209V	Page 24
S144P/N190F/Q209V	Page 25
E16P/S144P/N190F/Q209V	Page 25
M107L/Q167E/Y169K/N190F/Q209V	Page 26
N49S/M107L/Q167E/Y169K/N190F/Q209V	Page 26
N49S/M107L/H205R/Q167E/Y169K/N190F/Q209V	Page 26
LA-K38AMY	Pages 26-27
LA-K38AMY/Q167E/Y169K/N190F/Q209V	Page 27

These mutant $\alpha\text{-amylases}$ all have at least 95% sequence identity to SEQ ID NO:1.

As such, contrary to the Examiner's remarks, claim 1 does not encompass an unduly broad number of species. For this reason, the rejection is improper. Withdrawal thereof is respectfully requested.

Claims 5 and 6

Regarding claims 5 and 6, the Examiner appears to assert that the recitation of "two kinds of mutations" renders the

claims open to any number of mutations and represents an enormous genus of mutant α -amylases. In order to answer this rejection, claims 5 and 6 are amended to recite "a first mutation" and "a second mutation." Thus, the instant rejection is overcome.

Rejection under 35 U.S.C. § 112, first paragraph, enablement Claims 1-6, 10, and 11

The Examiner rejects claims 1-6, 10, and 11 under 35 U.S.C. § 112, first paragraph, for allegedly not being enabled by the specification. Claims 2, 4, and 11 are canceled, thus rendering rejection thereof moot. Applicants respectfully traverse the rejection applied to the pending claims. Reconsideration of the claims and withdrawal of the instant rejection are respectfully requested.

The Examiner maintains her assertion that the pending claims are not enabled by the specification. Specifically, the Examiner again asserts that claim 1 encompasses a protein variant having an untold number of mutations. Applicants respectfully disagree.

As noted above, the mutant α -amylase recited in claim 1 must (1) be derived from an amino acid sequence corresponding to SEQ ID NO:1 or an amino acid sequence 70% homologous to SEQ ID NO:1, (2) have a substitution or deletion of the 11th Tyr, 16th Glu, 49th Asn, 84th Glu, 144th Ser, 167th Gln, 169th Tyr, 178th Ala,

188th Glu, 190th Asn, 205th His, or the 209th Gln, (3) possess the specific functions of increased heat resistance and resistance to chelating agents, and (4) have a resulting amino acid sequence which is at least 70% homologous to SEQ ID NO:1. Thus, the mutant α -amylase mutant is defined by several limitations in the claim.

As such, contrary to the Examiner's remarks, the claims do not encompass an unduly broad number of species that are not enabled by the specification. For this reason, the rejection is improper. Withdrawal thereof is respectfully requested.

Rejection under 35 U.S.C. § 112, second paragraph

The Examiner rejects claims 1-6 and 10 under 35 U.S.C. § 112, second paragraph for allegedly being indefinite. Claims 2 and 4 are canceled, thus rendering rejection thereof moot. Applicants respectfully traverse the rejection applied to the pending claims. Reconsideration of the claims and withdrawal of the instant rejection are respectfully requested.

Claim 1

The Examiner asserts that claim 1 is unclear for reciting "at least one amino acid residue." Applicants respectfully disagree. Claim 1 is directed to a mutant α -amylase obtained by making a substitution or deletion of at least one amino acid

residue of specific positions in SEQ ID NO:1 or a homolog of SEQ ID NO:1. Applicants respectfully submit that it is clear from this language that at least one of the amino acid substitution/deletion must correspond to the 11th Tyr, 16th Glu, 49th Asn, 84th Glu, 144th Ser, 167th Gln, 169th Tyr, 178th Ala, 188th Glu, 190th Asn, 205th His, or the 209th Gln as noted in the claim. Thus, the rejection is improper and should be withdrawn.

Claims 2-4

The Examiner rejects claims 2-4 for reciting "liquefying α -amylase." Claims 2 and 4 are canceled. Regarding claim 3, the Examiner asserts the specification does not clearly define what is a "liquefying α -amylase." Claim 3, as amended, does not recite a "liquefying α -amylase." Thus, the instant rejection is overcome.

Claims 2 and 5

The Examiner rejects claims 2 and 5 for reciting "a substitution of a sequence...from the amino terminus...." Claim 2 is canceled. Regarding claim 5, the Examiner asserts that the exact position of the amino terminus is unclear. Applicants amend claim 5 to clarify that the second mutation consists of a substitution of a sequence corresponding to the 11th amino acid to the 100th amino acid from the N-terminal end of the sequence

(i.e., the first amino acid). Thus, the instant rejection is overcome.

Claim 5

The Examiner asserts that "a first (second) mutation is" in claim 5 is unclear. The phrase is amended to "a first (second) mutation consists of." Thus, the instant rejection is overcome.

Claim 6

Similarly, the Examiner asserts that "first (second) mutation comprises" in claim 6 is contradictory to claim 5, which recites "consists of." Claim 6 is amended to recite "a first (second) mutation consists of." Thus, the instant rejection is overcome.

Conclusion

Applicants respectfully submit that the above amendments and/or remarks fully address and overcome the rejections and objections of record. The instant claims are now in condition for allowance. Early and favorable action by the Examiner is respectfully requested.

If there are any minor matters precluding allowance of the application which may be resolved by a telephone discussion, the Examiner is respectfully requested to contact Kristi L. Rupert, Ph.D. (Reg. No. 45,702) at (703) 205-8000.

Pursuant to the provisions of 37 C.F.R. §§ 1.17 and 1.136(a), the Applicants hereby petition for an extension of two (2) months to July 25, 2003, in which to file a reply to the Office Action. The required fee of \$410.00 is enclosed herewith.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. § 1.16 or under 37 C.F.R. § 1.17; particularly, extension of time fees.

Respectfully submitted,

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Ву

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JWB/KLR Attachment:

Version with Markings to Show Changes Made Exhibits 1 and 2



VERSION WITH MARKINGS TO SHOW CHANGES MADE

In the Claims:

Claims 2, 4, and 11 are canceled.

Claims 15-27 are added.

The following claims are amended:

3. (Twice Amended) [The] A mutant α -amylase [according to Claim 2,] obtained by making a substitution of [wherein] an amino terminal sequence from 1st Asp through 19th Gly of SEQ ID NO:1 or an amino terminal sequence corresponding to 1st Asp through 19th Gly of SEQ ID NO:1 of a sequence having at least 95% [70%] homology to SEQ ID NO:1, [is substituted] with an amino acid sequence from 1st His to 21st Gly of SEQ ID NO:2, [encoding another liquefying α -amylase]

wherein said mutant α -amylase possesses increased heat resistance and maintains resistance to chelating agents when compared to SEQ ID NO:1.

5. (Three Times Amended) A mutant α -amylase obtained by introducing a first mutation and a second mutation [two kinds of mutations] into SEQ ID NO:1 or an amino acid sequence having at least 95% [70%] homology to SEQ ID NO:1,

wherein <u>said</u> [a] first mutation <u>consists of</u> [is] a substitution or a deletion of at least one amino acid residue selected from the group consisting of the 11th Tyr, 16th Glu, 49th Asn, 84th Glu, 144th Ser, 167th Gln, 169th Tyr, 178th Ala, 188th Glu, 190th Asn, 205th His and 209th Gln, and

wherein <u>said</u> [a] second mutation <u>consists of</u> [is] a substitution of a sequence corresponding to <u>the 11th to 100th</u> [11 to 100] amino acid <u>residue</u> [residues] from the amino terminus of the amino acid sequence set forth in SEQ ID NO:1, and

wherein said mutant α -amylase possesses increased heat resistance and maintains resistance to chelating agents when compared to SEQ ID NO:1.

6. (Twice Amended) The mutant α -amylase according to Claim 5, wherein said first mutation consists of [comprises]:

the substitution of an amino acid residue selected from the group consisting of: the 11th Tyr of SEQ ID NO:1 with Phe, the 16th Glu of SEQ ID NO:1 with Pro, the 49th Asn of SEQ ID NO:1 with Ser, the 167 Gln of SEQ ID NO:1 with Glu, the 169th Tyr of SEQ ID NO:1 with Lys, the 190th Asn of SEQ ID NO:1 with Phe, the 205th His of SEQ ID NO:1 with Arg, and the 209th Gln of SEQ ID NO:1 with Val,

and wherein said second mutation consists of [comprises]:

substituting an amino terminal sequence from $1^{\rm st}$ Asp through $19^{\rm th}$ Gly of SEQ ID NO:1 with an amino acid sequence from $1^{\rm st}$ His to $21^{\rm st}$ Gly of SEQ ID NO:2.

13. (Amended) A mutant α -amylase obtained by making a substitution or deletion of at least one amino acid residue of specific positions in SEQ ID NO:1, or by making a substitution or deletion of at least one amino acid residue corresponding to the above-mentioned amino acid residue in a sequence having at least 95% homology to SEQ ID NO:1,

wherein said at least one amino acid residue is selected from the group consisting of:

the 11^{th} Tyr, 16^{th} Glu, 49^{th} Asn, 84^{th} Glu, 144^{th} Ser, 167^{th} Gln, 169^{th} Tyr, 178^{th} Ala, 188^{th} Glu, 190^{th} Asn, 205^{th} His and 209^{th} Gln, and

wherein said mutant α -amylase:

- (i) decomposes $_{\alpha}\text{--}1,4\text{--glycoside}$ bonds of starch, amylose, amylopectin, and partially decomposed products thereof;
- (ii) produces glucose, maltose, maltotriose,
 maltotetraose, maltopentaose, maltohexaose, and
 maltoheptaose from amylose;
 - (iii) does not act on pullulan;

- (iv) exhibits a residual activity of at least 70% in a pH range of 6.5 to 11 under treatment conditions of 40° C and 30 minutes;
 - (v) acts in a temperature range of 20°C to 80°C;
- (vi) exhibits a residual activity of at least 80% when incubated at 40° C, or at least 60% when incubated at 45° C, for 30 minutes in 50 mM glycine-sodium hydroxide buffer at pH 10;
- (vii) has a molecular weight of $55,000 \pm 5,000$ as measured by sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis;
- (viii) has an isoelectric point of about 4.2 as measured by isoelectric focusing;
- (ix) has a residual activity of at least 90% when treated at pH 10 and 30°C for 30 minutes in a 0.1% solution of a surfactant selected from the group consisting of:

sodium linear alkylbenzenesulfonates, sodium alkylsulfates, sodium polyoxyethylene alkylsulfates, sodium $\alpha\text{-olefinsulfonates},$ sodium salts of $\alpha\text{-}$ sulfonated fatty acid esters, sodium alkylsulfonates, SDS, soap, and Softanol;

(x) is inhibited by 1 mM $\rm Mn^{2+}$ by about 75%, or by 1 mM $\rm Sr^{2+}$ or 1 mM $\rm Cd^{2+}$ by about 30 to 40%, when treated at pH 10 and 30°C for 30 minutes; and

(xii) comprises an amino acid sequence which is at least [95%] 70% homologous to SEQ ID NO:1.





SECOND SITE: http://www.ualberta.ca/~stothard/javascript/index.html The Sequence Manipulation Suite: Identity

SEQ ID NO 1 vs SEQ ID NO 4 Alignment length (not including identical gaps): 480

Identical residues: 463 Similar residues: 4 Percent identity: 96.5% Percent similarity: 97.3%

310

320

(A) 80 (B) 80 using	ison of: 655961 > SE 655962 > SE matrix file:	BLOSUM50, g			Ł		0 aa 5 aa
	10	20	30	40	50	60	
_	DGLNGTMMQYYEW	-					
	.: :::::::::::::::::::::::::::::::::::						
_	10	20	30	40	50	60	
	70	80	90	100	110	120	
_	LYDLGEFNQKGTV						
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-	LYDLGEFNQKGTV	RTKYGTRSQLÇ 80	90	31QVYGDVVMI 100	110	120	
	. •						
	130 NPTNRWODISGAY	140 TIDAWTGEDES	150 SCRNNAVSDEI	160 KWRWEHENGVI	170 WDORYO-EN	HTFRFAN	
_	: .:: :.::: :						
_	NRSNRNQEISGEY						
	130	140	150	160	170	180	
1	80 190	200	210	220	230		
_	TNWNWRVDEEN	GNYDYLLGSNI :::::::					
_	TGKAWDWEVDIEN						
	190	200	210	220	230	240	
	240 250	260	270	280	290		
_	FWYTSDWVRHQRN	~				-	
	. :: ::. : :: YSYTRDWLTHVRN						
_	250	260	270	280	290	300	
	300 310	320	330	340	350		
_	QGGSYDMRNILRG		/TFVDNHDTQI	PGESLESWVAI	OWFKPLAYAT	ILTREGG	
	.:: .:::::::::::::::::::::::::::::::::						
_	OGG T L DUKIN T DING	200	Y T. ADMIIDOQI	. CDADBOT VQC	VE DETUD	1111100	

330

340

350

360 370 380 390 400 410 YPNVFYGDYYGIPNDNISAKKDMIDELLDARQNYAYGTQHDYFDHWDVVGWTREGSSSRP YPSVFYGDYYGIPTHGVPSMKSKIDPLLQARQTYAYGTQHDYFDHHDIIGWTREGDSSHP 400 380 390 410 450 460 470 420 430 440 NSGLATIMSNGPGGSKWMYVGRQNAGQTWTDLTGNNGASVTINGDGWGEFFTNGGSVSVY NSGLATIMSDGPGGNKWMYVGKHKAGQVWRDITGNRSGTVTINADGWGNFTVNGGAVSVW 450 460 470 480 440 480 VNQ : . : VKQ

from: http://vega.igh.cnrs.fr/bin/lalign-guess.cgi

DATA USED:

>SEQ ID NO: 1

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>SEQ ID NO 4

DGLNGTMMQYYEWHLENDGQHWNRLHDDAEALSNAGITAIWIPPAYKGNSQADVGYGAYD LYDLGEFNQKGTVRTKYGTKAQLERAIGSLKSNDINVYGDVVMNHKLGADFTEAVQAVQV NPSNRWQDISGVYTIDAWTGFDFPGRNNAYSDFKWRWFHFNGVDWDQRYQENHLFRFANT NWNWRVDEENGNYDYLLGSNIDFSHPEVQEELKDWGSWFTDELDLDGYRLDAIKHIPFWY TSDWVRHQRSEADQDLFVVGEYWKDDVGALEFYLDEMNWEMSLFDVPLNYNFYRASKQGG SYDMRNILRGSLVEAHPIHAVTFVDNHDTQPGESLESWVADWFKPLAYATILTREGGYPN VFYGDYYGIPNDNISAKKDMIDELLDARQNYAYGTQHDYFDHWDIVGWTREGTSSRPNSG LATIMSNGPGGSKWMYVGQQHAGQTWTDLTGNHAASVTINGDGWGEFFTNGGSVSVYVNQ

>SEQ ID NO 2

HHNGTNGTMMQYFEWHLPNDGNHWNRLRDDAANLKSKGITAVWIPPAWKGTSQNDVGYGA YDLYDLGEFNQKGTVRTKYGTRSQLQGAVTSLKNNGIQVYGDVVMNHKGGADGTEMVNAV EVNRSNRNQEISGEYTIEAWTKFDFPGRGNTHSNFKWRWYHFDGTDWDQSRQLQNKIYKF RGTGKAWDWEVDIENGNYDYLMYADIDMDHPEVINELRNWGVWYTNTLNLDGFRIDAVKH IKYSYTRDWLTHVRNTTGKPMFAVAEFWKNDLAAIENYLNKTSWNHSVFDVPLHYNLYNA SNSGGYFDMRNILNGSVVQKHPIHAVTFVDNHDSQPGEALESFVQSWFKPLAYALILTRE QGYPSVFYGDYYGIPTHGVPSMKSKIDPLLQARQTYAYGTQHDYFDHHDIIGWTREGDSS HPNSGLATIMSDGPGGNKWMYVGKHKAGQVWRDITGNRSGTVTINADGWGNFTVNGGAVS VWVKQ



- 480 aa - 480 aa

Comparison of:

(A) 77810651 >_ SEQ ID NO: 1

(B) 77810652 >_ SEQ ID NO: 4

using matrix file: BLOSUM50, gap penalties: -14/-4

96.5% identity in 480 aa overlap; score: 3335

	10	20	30	40	50	60
	DGLNGTMMQYYEW	ILENDGQHWNR	LHDDAAALSI	DAGITAIWIPI	PAYKGNSQAD	VGYGAYD
_	:::::::::::::::::::::::::::::::::::::::	: : : : : : : : : :	::::: :::			::::::
	DGLNGTMMQYYEW	ILENDGOHWNR	LHDDAEALSI	NAGITAIWIPI	PAYKGNSOAD	VGYGAYD
_	10	20	30	40	50	60
	10	20	30	40	50	00
	70	80	90	100	110	120
_	LYDLGEFNQKGTV	RTKYGTKAQLE	RAIGSLKSN	DINVYGDVVMI	NHKMGADFTE.	AVQAVQV
	:::::::::::::::::::::::::::::::::::::::	: : : : : : : : : :	::::::::	:::::::::	:::::::	::::::
	LYDLGEFNQKGTV	RTKYGTKAOLE	RAIGSLKSN	DINVYGDVVM	NHKLGADFTE.	AVQAVQV
_	70	80	90	100	110	120
	, •	00	30			
	120	140	150	160	170	180
	130	140				
_	NPTNRWQDISGAY					
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	NPSNRWQDISGVY	ridawtgfdff	GRNNAYSDF	KWRWFHFNGVI	DWDQRYQENH	LFRFANT
_	130	140	150	160	170	180
	190	200	210	220	230	240
	NWNWRVDEENGNYI					
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_	NWNWRVDEENGNY	DYLLGSNIDFS	HPEVQEELK	DWGSWFTDELI	OLDGYRLDAI	KHIPFWY
	190	200	210	220	230	240
	250	260	270	280	290	300
	TCDWADAODNEAD	NT.EVAGEVWK	יחחעמאַז.בּבּעי	LIPMNWEMSI.I	TOVDI.NVNFV	RASOOGG
_	TSDWVRHQRNEAD	_				
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-	TSDWVRHQRSEAD	DLFVVGEYWK	::::::: DDVGALEFY	:::::::: LDEMNWEMSL	FDVPLNYNFY	::::: RASKQGG
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_	TSDWVRHQRSEAD	DLFVVGEYWK	::::::: DDVGALEFY	:::::::: LDEMNWEMSL	FDVPLNYNFY	::::: RASKQGG
- -	TSDWVRHQRSEAD	DLFVVGEYWK	::::::: DDVGALEFY	:::::::: LDEMNWEMSL	FDVPLNYNFY	::::: RASKQGG
_	TSDWVRHQRSEAD(250	260 320	::::::: DDVGALEFY: 270	::::::::::::::::::::::::::::::::::::::	:::::::: FDVPLNYNFY 290 350	:::.:: RASKQGG 300
- -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI	EIIIIIIIII QDLFVVGEYWK 260 320 EAHPMHAVTFV	DDVGALEFY 270 330 DNHDTQPGE	::::::: LDEMNWEMSLI 280 340 SLESWVADWFI	:::::::: FDVPLNYNFY 290 350 KPLAYATILT	:::.:: RASKQGG 300 360 REGGYPN
- -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI	2001 2001 200 320 200 200 200 201 201 201 201 201 201 2	DDVGALEFY 270 330 DNHDTQPGE	:::::::: LDEMNWEMSL1 280 340 SLESWVADWF1	:::::::: FDVPLNYNFY 290 350 KPLAYATILT	:::.:: RASKQGG 300 360 REGGYPN ::::::
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI	ELLER STEEL	DDVGALEFY 270 330 DNHDTQPGE CONHDTQPGE	:::::::: LDEMNWEMSLI 280 340 SLESWVADWFI ::::::::	:::::::: FDVPLNYNFY 290 350 KPLAYATILT ::::::::	:::.:: RASKQGG 300 360 REGGYPN ::::::
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI	2001 2001 200 320 200 200 200 201 201 201 201 201 201 2	DDVGALEFY 270 330 DNHDTQPGE	:::::::: LDEMNWEMSL1 280 340 SLESWVADWF1	:::::::: FDVPLNYNFY 290 350 KPLAYATILT	:::.:: RASKQGG 300 360 REGGYPN ::::::
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI ::::::::::::::::::::::::::::::::::::	E:::::::: QDLFVVGEYWK 260 320 EAHPMHAVTFV EAHPIHAVTFV 320	DDVGALEFY 270 330 DNHDTQPGE CONHDTQPGE 330	::::::::: LDEMNWEMSLI 280 340 SLESWVADWFI ::::::::: SLESWVADWFI 340	::::::::::::::::::::::::::::::::::::::	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI SYDMRNILRGSLVI 310 370	200 320 320 EAHPMHAVTFV 320 320 380	270 330 DNHDTQPGE: :::::::::::::::::::::::::::::::::::	::::::::::::::::::::::::::::::::::::::	::::::::::::::::::::::::::::::::::::::	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI ::::::::::::::::::::::::::::::::::::	200 320 320 EAHPMHAVTFV 320 320 380	270 330 DNHDTQPGE: :::::::::::::::::::::::::::::::::::	::::::::::::::::::::::::::::::::::::::	::::::::::::::::::::::::::::::::::::::	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI SYDMRNILRGSLVI 310 370	260 320 320 EAHPMHAVTFV 320 380 ISAKKDMIDEL	DDVGALEFY 270 330 DNHDTQPGE CONHDTQPGE 330 390 LDARQNYAY	::::::::::::::::::::::::::::::::::::::	::::::::::::::::::::::::::::::::::::::	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI	260 320 EAHPMHAVTFV 320 EAHPIHAVTFV 320 380 ISAKKDMIDEL	DDVGALEFY 270 330 DNHDTQPGE 330 DNHDTQPGE 330 390 LDARQNYAYO	::::::::::::::::::::::::::::::::::::::	FDVPLNYNFY 290 350 KPLAYATILT CPLAYATILT 350 410 DVVGWTREGS	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG ::::::
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI ::::::::::::::::::::::::::::::::::::	260 320 EAHPMHAVTFV 320 EAHPIHAVTFV 320 380 ISAKKDMIDEL	DDVGALEFY 270 330 DNHDTQPGE 330 390 LDARQNYAY LDARQNYAY	::::::::: LDEMNWEMSLI 280 340 SLESWVADWFI 340 400 GTQHDYFDHWI :::::::::	FDVPLNYNFY 290 350 KPLAYATILT STORMART	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG ::::::: SSRPNSG
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI	260 320 EAHPMHAVTFV 320 EAHPIHAVTFV 320 380 ISAKKDMIDEL	DDVGALEFY 270 330 DNHDTQPGE 330 DNHDTQPGE 330 390 LDARQNYAYO	::::::::::::::::::::::::::::::::::::::	FDVPLNYNFY 290 350 KPLAYATILT CPLAYATILT 350 410 DVVGWTREGS	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG ::::::
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI VFYGDYYGIPNDNI 370	260 320 EAHPMHAVTFV 320 380 ISAKKDMIDEL 380 ISAKKDMIDEL 380	DDVGALEFY 270 330 DNHDTQPGE 330 390 LDARQNYAY 390	::::::::::::::::::::::::::::::::::::::	FDVPLNYNFY 290 350 KPLAYATILT STORMARY 350 410 DVVGWTREGS STORMARY 410	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG ::::::: SSRPNSG 420
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI VFYGDYYGIPNDNI 370 430	260 320 2AHPMHAVTFV 320 380 ISAKKDMIDEL 380 440	270 330 DNHDTQPGE: 330 390 LDARQNYAY0 390 450	::::::::::::::::::::::::::::::::::::::	### ##################################	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG ::::::: SSRPNSG 420 480
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI VFYGDYYGIPNDNI 370	260 320 2AHPMHAVTFV 320 380 ISAKKDMIDEL 380 440	270 330 DNHDTQPGE: 330 390 LDARQNYAY0 390 450	::::::::::::::::::::::::::::::::::::::	### ##################################	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG ::::::: SSRPNSG 420 480
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI VFYGDYYGIPNDNI 370 430	260 320 2AHPMHAVTFV 320 380 ISAKKDMIDEL 380 440 WMYVGRQNAGQ	DDVGALEFY 270 330 DNHDTQPGE 330 390 LDARQNYAY 390 LDARQNYAY 390 450	::::::::::::::::::::::::::::::::::::::	FDVPLNYNFY 290 350 KPLAYATILT STON A 10 DVVGWTREGS COLVGWTREGT 410 470 NGEFFTNGGS	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG :::::: SSRPNSG 420 480 VSVYVNQ
- - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI 370 VFYGDYYGIPNDNI 370 430 LATIMSNGPGGSKI	260 320 2AHPMHAVTFV 320 380 ISAKKDMIDEL 380 ISAKKDMIDEL 380 440 WMYVGRQNAGQ	DDVGALEFY 270 330 DNHDTQPGE 330 390 LDARQNYAY 390 450 TTWTDLTGNNO CHARTONYON CH	::::::::::::::::::::::::::::::::::::::	FDVPLNYNFY 290 350 KPLAYATILT STORMAR AND	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG :::::: SSRPNSG 420 480 VSVYVNQ ::::::
- - - -	TSDWVRHQRSEADO 250 310 SYDMRNILRGSLVI 310 370 VFYGDYYGIPNDNI 370 VFYGDYYGIPNDNI 370 430 LATIMSNGPGGSKI	260 320 2AHPMHAVTFV 320 380 ISAKKDMIDEL 380 ISAKKDMIDEL 380 440 WMYVGRQNAGQ	DDVGALEFY 270 330 DNHDTQPGE 330 390 LDARQNYAY 390 450 TTWTDLTGNNO CHARTONYON CH	::::::::::::::::::::::::::::::::::::::	FDVPLNYNFY 290 350 KPLAYATILT STORMAR AND	:::.:: RASKQGG 300 360 REGGYPN :::::: REGGYPN 360 420 SSRPNSG :::::: SSRPNSG 420 480 VSVYVNQ ::::::

done at: http://vega.igh.cnrs.fr/bin/lalign-guess.cgi